Orthogonal Recursive Bisection on the GPU for Accelerated Load Balancing in Large N-Body Simulations

Bachelor Thesis

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Contents

1	Introduction					
2	Bac	Background				
	2.1	Fast Multipole Expansion	5			
		2.1.1 Advantage of Cubic Bounding Volumes	6			
	2.2	ORB vs. K-d Trees	6			
	2.3	Rise of the GPU	6			
	2.4	Related Work	7			
3	Ort	hogonal Recursive Bisection (ORB)	9			
	3.1	Target Data and Notation	9			
	3.2	Memory and Workload Balancing	10			
	3.3	ORB	11			
		3.3.1 Algorithm	12			
	3.4	By Example	13			
	3.5	Root finding	16			
		3.5.1 Bisection Method	17			
		3.5.2 Edge Cases	18			
		3.5.3 Runtime Analysis	19			
	3.6	Partition Algorithm	19			
4	The	eoretical Analysis	21			
	4.1	General Memory Model	21			
	4.2	Supercomputers	22			
	4.3	Roofline Performance Model	23			
		4.3.1 Estimating Flops	23			
		4.3.2 Estimating Arithmetic Intensity	24			
		4.3.3 The Plots	25			
		4.3.4 Empirical Verification	28			
	4.4	Runtime Estimates	29			
		4.4.1 CPU Version	29			
		4.4.2 GPU Counting	30			
		4.4.3 GPU Counting and Partitioning	30			
		4.4.4 Plugin Values	30			
	4.5	Conclusion	31			
5	\mathbf{CP}^{T}	U Implementation	33			
	5.1	MDL and PKDGrav	33			
	5.2	Particles	33			
	5.3	Cell	34			
		5.3.1 Heap Conditions	34			
		5.3.2 Class	35			
	5.4	Mapping Cells to Particles	36			
	5.5	Services	36			

		5.5.1 Init and Finalize Service	36
		5.5.2 Count Left Service	36
		5.5.3 Count	37
		5.5.4 Partition	38
		5.5.5 Make Axis	39
	5.6	Parallel Schedule	39
6	GP	U Implementation	42
	6.1	Relevant CUDA Concepts	42
			42
		6.1.2 Memory	43
		6.1.3 Memory Access Patterns	43
		6.1.4 Asynchronous Operations	44
		6.1.5 Synchronization	44
	6.2		45
	6.3	Memory Management	45
	6.4	GPU Accelerated Count Left	45
		6.4.1 Schedule	45
		6.4.2 Service	47
		6.4.3 Kernel Code	48
	6.5	Improved GPU Accelerated Count Left	50
		6.5.1 Schedule	51
			52
	6.6	GPU Accelerated Partitioning	55
		6.6.1 Schedule	56
		6.6.2 Partitioning a Single Cell	57
			57
7	Per	v	59
	7.1	Scaling	60
8	Cor	nclusion	61

1 Introduction

The N-Body technique has been used for decades to simulate the Universe to compare theory with observations. It uses "particles" to represent objects of a certain mass and computes the forces by applying the gravity equation. As the forces operate over infinite distance it is necessary to consider all pair-wise interactions, making a naive implementation $\mathcal{O}(n^2)$. This does not scale with large particle counts. (doug) the problem becomes "computationally infeasible" with large particle counts

A common solution is to partition the space in which the particles are contained, into a set of subspaces, also called cells. These cells are then stored in a space partitioning tree data structure (SPTDS) in order to speed up the simulation with the Fast Multipole Method (FMM) to O(n). Building the data structure uses a significant percentage of the overall simulation time. As of now this was usually done on the CPU, not leveraging GPU acceleration. In this thesis an upper limit for the speedup of GPU accelerated SPTDS building algorithms over its CPU only counterpart is established. An efficient methods of building the SPTDS on the CPU and the GPU is then proposed and implemented in C++ and CUDA.

TODO: Describe measurement results here

2 Background

Structures occurring within astrophysical simulations can be identified (doug) we model observations (create mock universes) to compare with observation in real observed data [Stadel, 2001]. Thus the simulations can help us understand the universe. Generated datasets with similar distributions to reality are used to seed the simulations. Mass field distribution are approximated with a large number of particles, where each of them represents a mass point. (doug) We model a collisionless fluid and use "particles" to sample the mass distribution Numerical integration methods then compute the gravitational forces on each object. Originally developed by Joachim Stadel during his PhD at University of Washington [Stadel, 2001], PKDGrav is a fully parallel N-Body code which is now maintained and improved by the ICS at UHZ [UZH, 2022]. PkDGrav, as well as most state of the art simulation softwares, use a space partitioning tree data structure (SPTDS), which is used to partition a space into subspaces. Each cell (node), in the tree represents a subspace, and the children of the node represent the subspaces that are contained within the node's subspace. The mutlipole expansion is then applied to each cell of the SPTDS and the fast multipole method increases the runtime from $O(N^2)$ to O(N) where N is the total number of particles. [Stadel, 2001]

2.1 Fast Multipole Expansion

In mathematics, a multipole is a concept used to approximate the shape of a physical body (doug) distribution. In this case the body consists of the particles contained within the subspace, or volume of a cell. The multipole expansion is the mathematical series that results from this approximation. The resulting series can simplify the original body, by choosing a suitable precision.

To keep force integration within a reasonable error margin, in most cases it is not necessary to compute all N to N interactions. The fast multipole method can compute gravity in O(N) runtime by leveraging the SPTDS. [Stadel, 2001]

On a more intuitive level, one can imagine a large cluster of objects far away from planet Earth. To compute the interactions between every single object of the cluster and Earth, $O(N^2)$ have to be computed computations. However, since the object is reasonably far away, it does not make sense to compute every interaction. Instead, the cluster can be summarized into a single object and compute the gravity acting between planet Earth and the group. Consequently, objects within our own solar system would be too close to planet Earth to approximate its effects using a multipole. In such cases, it is necessary to perform all computations, however they can be hidden within a constant and do not affect the total runtime. Note that this example does not directly translate to the simulations, as a particle is not equivalent to a planet or a star. More on that in section 3.1.

2.1.1 Advantage of Cubic Bounding Volumes

The distance between the center of mass of a cell and its most distant corner correlate with the error ratio [Stadel, 2001], which in turn influences how many timesteps have to be performed. Of the family of rectangular cuboids, the cube is the shape where the average distance between any point and its most distant corner point is the smallest.

2.2 ORB vs. K-d Trees

A number of methods exist to create SPTDS, however since each cell on the same depth of the tree is required to encompass an almost equal number of particles, methods like octrees are not suitable as they do not produce balanced groups. The k-d tree construction method however is strikingly similar to the ORB method, however there is a slight difference in the construction process. The Orthogonal Recursive Bisection method constructs a SPTDS where all resulting cells represent subspaces, or volumes, similar to cubes. K-d construction methods generate SPTDS as well, but shapes of the resulting subspaces can be strongly skewed and are thus ill suited for FMM. Both algorithms start with a single cell and iteratively construct the SPTDS by searching for axis aligned planes cutting the volume of each leaf cell of the current SPTDS into two. From the resulting two volumes, two new cells are generated and appended to the SPTDS.

In k-d tree construction, the axis lying orthogonal to the cut plane is chosen periodically, meaning all cells with an equal distance (depth) to the root of the SPTDS are cut with planes orthogonal to the same axis. In the case of ORB, the axis where a cells volume is largest is picked to compute an orthogonal plane. Therefore cells with the same depth can have cut planes of variable orientations. This make ORB special as it combines a balanced distribution with near cubic shapes.

2.3 Rise of the GPU

Most if not all state of the art supercomputers are hybrid, meaning each node has a CPU as well as a GPU[TOP500, 2022]. GPU accelerated code can increase the speed of many computations, where in most cases CUDA is used to fully leverage the hardware.

The computational effort required in PKDGrav and most modern astrophysical N-Body simulations can be divided into three categories:

1. Global Tree Building / Load Balancing: In supercomputers, particles are distributed equally (doug) not "equally": to balance work or memory or some combination among computing nodes to leverage node level parallelization. ORB is used to generate a well suited SPTDS for FMM on a node level.

- 2. Local Tree Building: The same approach is repeated on a local level, where instead of distributing the workload among nodes, its distributed among threads. The tree from step 1. is expanded by the locally computed tree. (doug) tree building continues down to "buckets"
- 3. Force Calculation and Integration: The STPDS constructed in step 1. and 2. combined with FMM is used to compute force integration.

In PKDG each category is about one third of the total calculation time [?]. Making Global (1.) and Local Tree Building (2.) subjects to possible performance improvements, as GPU acceleration is as of now only exploited in Force Calculation and Integration(3.). (doug) actually just the force calculations

The main objective is to improve the runtime of astrophysical simulations, or more specifically PKDGrav. Simultaneously penalties with regards to N, the total number of particles, should be kept as low as possible. Simulation accuracy benefits from large N as well as an increased number of time steps. As GPU accelerations are low level, hardware specific details are relevant. Thus the algorithms should be adapted and optimized the algorithms with regards to the target hardware Piz Daint.

2.4 Related Work

A number of GPU accelerated k-d tree construction algorithms exists, however the methods cannot simply be translated to ORB due to different axis choosing method. Fore example in the master thesis of Voglsam [Voglsam, 2013] a k-d tree is computed on the GPU and used to improve ray tracing for 3D graphics. However it makes use of a binning algorithm to find an optimal cut across cells of the same depth, which can be done in a single iteration. However in this case the axes for cells of the same depth vary.

When subdividing ORB into its individual components as shown below, some subroutines can be computed using fairly general algorithms, which have already been implemented and optimized for the GPU in many open source libraries.

- 1. Cut cells on last level in tree
 - (a) Make cut plane position guess
 - (b) Count number of particles left to the cut plane
 - (c) Repeat 1. till correct plane was found
- 2. Partition particles
- 3. Repeat till desired depth was reached

Thrust by NVIDIA [NVIDIA, 2022f] has an implementation of a reduction, which can be used to perform step 1.b), a binary search is equivalent to the

entire step 1. and finally it also exposes a partition interface. The library is very high level and does not allow us to control memory operations, which are very crucial to a highly efficient implementation of ORB. Furthermore CUB [NVIDIA, 2022b], also developed by NVIDIA, exposes a more low level API of the equivalent elements but the implementation is not general enough to deal with some critical performance issues. As the SPTDS grows, so do the number of leaf cells and thus the number of cells for which a cut plane has to be found simultaneously. When initializing a kernel for each cell, the number of kernels initialized grow exponentially, resulting in kernel invocation overheads dominating the actual computational costs. The problem is almost equivalent to a Device Segmented Reduce[NVIDIA, 2022c] where CUB provides an API, however unfortunately the reduction operator is set to be the same across all segments, which is not true in this case as for each cell the particle coordinates are compared to a different value.

3 Orthogonal Recursive Bisection (ORB)

Each cell in the SPTDS is enriched with the following properties:

- V_{cell} 3D volume where all corresponding particles are contained within.
- d_{cell} Number of leaf cells to be found among all subsequent cells.
- N_{cell} Number of particles encompasses in the volume of the cell.

Naturally each cell has two pointers pointing to its two children, which are used to traverse the tree, parent cell pointers important but could be added as well.

3.1 Target Data and Notation

A modified random Gaussian distributed is used to generate the target dataset, where its set to fit measured constraints of the actual universe. Thus the number of particles can set dynamically. In the simulation, a single particle has a mass of 10⁸ solar masses. (doug) different simulations will use dramatically different particle masses. Are you giving an example here or making a more general statement?

- The space space of binary numbers with a precision of p is defined as \mathbb{B}_p where $a \in \mathbb{B}_p \Leftrightarrow a \in \{0,1\}^p$
- The corner coordinates of the root volume are defined as $lower, up\vec{p}er$ where $lower, up\vec{p}er \in \mathbb{B}_p^3$.
- The coordinates of a particle i are defined as $\vec{x_i}$ for which holds $\{\vec{x}|lower \leq \vec{x} \leq up\vec{p}er, \vec{x} \in \mathbb{B}_p^3\}$. When referring to a single coordinate of a particle object, it is done by writing $\vec{x}_{i,j}$. Finally the array of all particles positions in a single axis is denoted as $\vec{x}_{:,j}$.

It lies in the nature of the universe, that large clusters of particles are found in some places, whereas other areas may be vastly empty of any objects. This is a very crucial characterization and differentiates this dataset from many other application of FMM, where the distribution are more uniform.

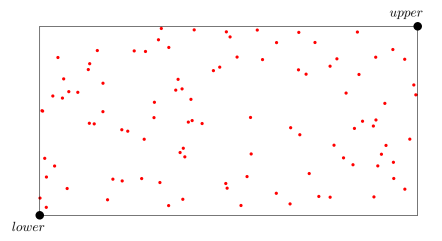


Figure 1: Uniform random distribution of 3D coordinates in cube domain projected onto a 2d plane

3.2 Memory and Workload Balancing

As mentioned the aim is to (1) improve the runtime of ORB whilst (2) at the same time keeping the number of particles N as large as possible. (doug) really? Don't we want to improve performance for any given N? There exists some averse effects when disregarding interactions between minimizing runtime and maximizing N.

To reduce the force integration error across all particles, some particles require more timesteps than others. This is caused by the vastly variant forces which are exerted on the particles. Whilst some follow a straight path with an almost constant velocity requiring little computational effort for great precision, others are influenced by strong gravitational poles resulting in highly curved movement paths and correlating with higher numbers of timesteps required.

The workload for a particle p_i is defined as the weighting function $w(p_i)$. The workload correlates with the number of simulations steps, that need to be computed for a single particle. In an optimally parallelized system balanced in terms of computational effort, the workload should be very similar among computing units. Where ideally the sum over all the particles considered by a thread is equal to the sum of all others. With strongly varying weights among particles, perfectly balancing the computational effort, results in a unbalanced distribution in terms of memory.

Lets consider a simple example (doug) this example could be simpler to illustrate the point. Given the set of particles $A = \{p_1, p_2, ..., p_{2m/3}\}$ and respectively $B = \{p_{2m/3+1}, p_2, ..., p_m\}$. This yields a total number of particles equivalent to N = m. Assuming that $\forall p \in A : w(p) = 1$ and $\forall p \in B : w(p) = 2$. All particles from set A are assigned to processor rank 0 and the particles from

B to rank 1. It follows $\sum_{p \in A} w(p) = \sum_{p \in B} w(p)$. Thus the two processors are balanced in terms of computing costs, but not in terms of memory size. In fact, process with rank 0 has 2m/3 elements and rank 1 has m/3 elements. Assuming each process has a memory size of 2m/3, then clearly this configuration is not optimal. When favoring memory balancing, 2m/3 could be assigned to each processor and $(4/3) \times m$ as many particles could be processed in total, which is larger than the original N and therefore favorable.

To which degree its ideal favor memory over workload balancing is difficult to answer. For now parametrize the workload using the weighting function. The workload balancing can also be completely ignored by setting $w(p_i) = 1$ for all particles.

3.3 ORB

To introduce the ORB algorithm, a recursive implementation of ORB is considered as it is easier to understand than the iterative version. All cell properties are defined recursively.

As the recursion is initialized with the root cell alone, it follows that V_{root} is equivalent to entire volume of the input data. $d_{cell} = d$ and $N_{root} = N$ are true per definition of N and d. When a cell is cut into two child cells, the children are refereed to as leftCell and rightCell. For any cell the following equations hold:

$$depth_{cell} = \lceil \log_2 d_{cell} \rceil \tag{1}$$

$$V_{cell} = V_{leftCell} \cup V_{rightCell} \tag{2}$$

$$V_{leftCell} \cap V_{rightCell} = \emptyset \tag{3}$$

Defining d_{cell} for each cell is especially important, since its crucial to ensure the final tree is a nearly complete binary tree. The way $d_{leftCell}$ and $d_{rightCell}$ is defined, guarantees that on the last level of tree the cells are filled from the left to the right. This way no gaps are present and the tree can be stored as a heap in an array.

$$d_{leftCell} = \min\{d_{cell} - 2^{depth_{cell} - 2}, 2^{depth_{cell} - 1}\}$$

$$\tag{4}$$

$$d_{rightCell} = \max\{d_{cell} - 2^{depth_{cell} - 1}, 2^{depth_{cell} - 2}\}$$
 (5)

Finally the number of particles encompassed in $V_{leftCell}$ and $V_{rigtCell}$ are defined as follows:

$$N_{leftCell} = min \left\{ x \in \{0, ..., N_{cell}\} : \sum_{i=0}^{x} w(p_i) \ge \frac{d_{leftCell}}{d_{cell}} \times \sum_{i=0}^{N_{cell}} w(p_i) \right\} - 1$$
(6)

$$N_{rightCell} = N_{cell} - N_{leftCell} \tag{7}$$

To simplify the visual and numeric explanation of the ORB algorithm, its assumed assume $\forall i \in \{0,..,N\} : w(p_i) = 1$.

The cut plane, which divides $V_{leftCell}$ and $V_{rightCell}$ is axis aligned an thus orthogonal to a given axis a. Thus searching for a single value c is sufficient to construct the plane, where c is the position of the plane along the axis a. The plane is considered ideal if the number of particles where $x_{i,a} \leq c$ are equivalent to $N_{leftCell}$.

After a plane is found, the V_{cell} can be divided or cut in two volumes, where $V_{leftCell}$ and $V_{rightCell}$ are constrained by the original V_{cell} and the cut plane. As defined in equation 3 the volumes do not intersect, thus $V_{leftCell}$ along axis a ends at c, where $V_{rightCell}$ start at c.

3.3.1 Algorithm

Algorithm 1 as mentioned below, is the main routine of ORB. All the volumes are described using a lower and an upper boundary point, sufficiently describing a box, thus the only input parameters required is x storing the particles positions, lower, upper and finally d. If d is equal to 1, this means the reduction must not be continued, as the target of a single subsequent leaf cell is already reached, since the cell itself is treated as a leaf cell. The stopping condition is formulated on lines 2-4. Line 6 describes a call to a method called maxIndex(), which essentially compares all provided values and returns the index of the maximum values. The result i provides us with the axis where the cell volume is largest. $d_{leftCell}$ is computed as described in equation 4 and its results is stored in a variable named d'. All prerequisites are met to compute the actual cut with the cut method which will be explained in more detail in section 3.5. The return value stored in cut is equivalent to the position of the cut plane along the cut axis. Using the cut value, the array of particles is partitioned as described in 3.6 returning the mid value which is the pivot index of the partitioning. Finally the original volume can be subdivided into two volumes as described on lines 10-13 and recursively call ORB method passing it two slices of the x array. **TODO:** Adapt lines numbers

Algorithm 1 The ORB main routine

```
1: procedure ORB(x, lower, up\vec{p}er, d)
         if d = 1 then
 2:
             return
                                                                             3:
         end if
 4:
         \vec{size} = u\vec{pper} - \vec{lower}
 5:
         i = maxIndex(\vec{size_0}, \vec{size_1}, \vec{size_2})
                                                                      ▷ Get index of max value
 6:
 7:
        l = \lceil \log_2 d \rceil
                                                                                             ▷ Depth
         d' = \min^{2} \{d - 2^{l-2}, 2^{l-1}\}
 8:
         cut = cut(x, \overrightarrow{lower_i}, u\overrightarrow{pper_i}, i, \frac{d'}{d})
                                                                                  ▶ Find cut plane
 9:
         mid = partition(x, split, axis)
                                                                             ▶ Partition particles
10:
         up\vec{p}er' = up\vec{p}er
11:
         upperChild_i = cut
12:
         \vec{lower'} = \vec{lower}
13:
         lowerChild_i = cut
14:
         ORB(x_{0:mid,:}, lower, upper', d')
15:
         ORB(x_{mid:len(x),:} \overrightarrow{lower'}, \overrightarrow{upper}, d-d')
16:
17: end procedure
```

3.4 By Example

A sample dataset is proposed to help with visual and numerical explanations of the algorithm. All particles in the sample dataset are assumed to have a weight of 1.

The algorithm is explained visually using the following example data set:

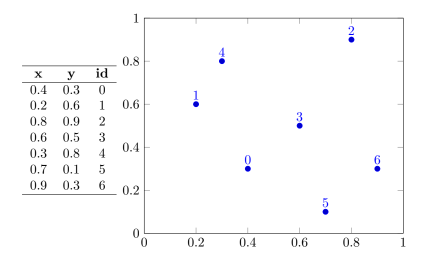


Figure 2: Example distribution with N=7

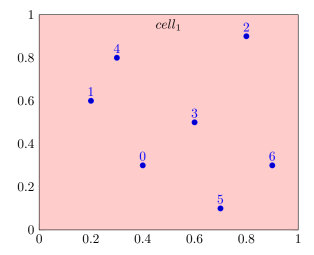


Figure 3: Example particles with ORB at recursion depth 0



Figure 4: Tree with ORB at recursion depth 0

Depicted in figure 3 & 4 is the SPTDS after initialization: there is only one

 $cell_1$ which is root and V_{cell_1} encompasses the entire domain which in this case is the rectangle ranging from 0 to 1 in both the x and y axis. d_{cell_1} is equivalent to 3 and N_{cell_1} is 7.

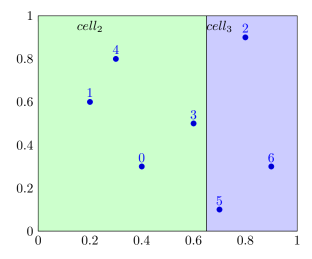


Figure 5: Example particles with ORB at recursion depth 1



Figure 6: Tree with ORB at recursion depth 1

A cut plane, which translates to a cut line in the simplified 2D example is constructed and $cell_2$ and $cell_3$ are generated accordingly.

5 & 6 Depict a SPTDS of depth 2, or two levels, where a cut plane was found dividing V_{cell_1} into V_{cell_2} and V_{cell_3} , $d_{cell_2}=2$ and $d_{cell_3}=1$ subsequently are calculated, meaning $cell_3$ is considered a leaf cell. Thus N_{cell_2} can be computed using a simplified version of equation 6 since all weights are assumed to be 1: $N_{cell_2} = \lfloor \frac{d_{cell_2}}{d_{cell_3}} * N_{cell_1} \rfloor = \lfloor \frac{2}{3} * 7 \rfloor = 4$ subsequently $N_{cell_3}=3$.

Recursion depth 2

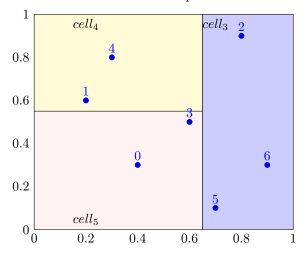


Figure 7: Example particles with ORB at recursion depth 2

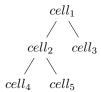


Figure 8: Tree with ORB at recursion depth 2

Again the procedure is repeated for $cell_2$ but not for $cell_3$ as there recursion is terminated by the stopping condition. Finally $cell_4$ and $cell_5$ is computed, where the stopping condition is met as well. The resulting SPTDS has 3 leaf cells and partitions the space into three subspaces.

3.5 Root finding

The cut algorithm takes an array of particle positions x, an axis, left and right boundaries and a percentage. Its goal is to return a position along the cut axis such that the particles less or equal to the a cut value are equivalent to the percentage multiplied by the length of the x array.

The problem is related to a selection algorithm and a quick select [Cormen et al., 2022] could be used. However quick select has a worst case runtime of $O(n^2)$. Furthermore median algorithms could be explored as well, but they are mostly approximation algorithms with bad worst case runtimes. The problem can be reformulated as a root finding problem. To do so a function f(c) is defined

which evaluates the number of elements smaller than the cut value minus half (or any other desired percentage) the number of total particles. When the function evaluates to zero, a an axis aligned cut plane is found, where exactly half the particles are located on the left side. There exist many different solvers for the root-finding problem, but the most stable and easiest to implement is the bisection method. Some solver combine approximate solver with the stable bisection method to generate fast but stable root finding methods. Such algorithms could be explored in later work.

Since the algorithm operates on binary numbers with a limited precision, the bisection method is guaranteed to terminate after p steps.

3.5.1 Bisection Method

Initially an estimation for a cut is made, in this case the exact middle of the domain boundaries. Some median finding algorithms use improved guessing to speed up the process, again a method which could be leveraged in later work.

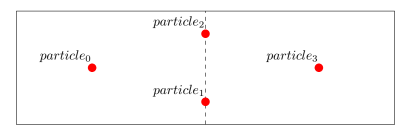
Because the maximal number of iterations is known, a loop can be used as seen below in algorithm 2 on line 4. A cut is then computed (line 5), which in the first iteration is the center position between left and right. It is then checked whether the stopping condition has already been reached (line 7-9), meaning the cut lies within α points of the ideal result. If it was not reached, the boundaries can be improved as follows: In case there are too many elements left of the cut, it is known that the cut position was chosen too far to the right and it follows that the ideal cut must be left of the current guess. Therefore, the boundaries can be adjusted, in this case by setting righ equal to cut as seen on line 13. The analogous concept can be applied in the other case (line 11).

Algorithm 2 Bisection Method

```
1: procedure CUT(x, left, right, axis, percentage)
2:
       nLeft = 0
3:
       cut = 0
       for k \in 0,..,p do
4:
          cut = (right + left)/2
5:
          nLeft \leftarrow sum(x_{:,axis} < cut)
                                                ▷ Counting particles left of cut
6:
7:
          if abs(nLeft - len(x)) * percentage < \alpha then
             Break
                                                          8:
          end if
9:
          if nLeft \le len(x) * percentage then
10:
11:
             left = cut
          else
12:
              right = cut
13:
          end if
14:
       end for
15:
16:
       return cut
17: end procedure
```

3.5.2 Edge Cases

Considering the following example particle distribution where $particle_1$ and $particle_2$ have identical x coordinates.



In this case there exists no ideal cut in the x (horizontal) axis. As either there is 1 particle to the left or 3, but no cut can result in 2 particles to the left. If more particles are added along the same line, the method performs even worse. However since the algorithm uses a deterministic for loop, the algorithm will terminate anyways and puts either 3 particles to the left or the right.

3.5.3 Runtime Analysis

On line 5, the number of particles to the left of the cut plane are summed. As the array is unordered, all coordinates of one axis need to be read once, resulting in a runtime of O(N). All other operations inside the for loop can be computed in a negligible constant time. The loop itself is repeated p number of times, where p corresponds to the precision of x coordinates.

To proof that the loop concludes after p iterations for integer numbers. With each iteration the range of possible solutions is divided by two. The same goes for integer numbers, each time a bit is removed, the size of the range of numbers which can be represented is reduced by two. Thus after p iterations the precision limit is reached and the cut cannot be improved. The proof does not hold analogously for float numbers as the precision is uniformly distributed there, however its also limited.

3.6 Partition Algorithm

Its desired to continuously update the array storing the positions of the particles and in the words used before, have a direct correlation between x_i and i. This enables grouping particles within a cell in a fixed range of two indexes of the particles array. The advantages of this are two fold: Access all particles within a cell in constant time, manipulate particles within a cell using a slice of the array.

As seen in the pseudo code in algorithm 3, the algorithm looks for a pair of particles, where for both the coordinate along the relevant axis are on the wrong side of the cut plane. In this case the particles can be swapped (line 8) resulting in the correct position of both particles. Correctness and a more detailed explanation of the algorithm can be found in the book [Cormen et al., 2022]. **TODO:** citations

Algorithm 3 Partition Method

```
1: procedure PARTITION(x, cut, axis)
        i = 0
 2:
 3:
        for k \in 0, ...N - 1 do
            if \vec{x}_{k,axis} \leq cut then
 4:
                while \vec{x}_{i,axis} \leq cut and i < N do
 5:
                    i = i + 1
 6:
                end while
 7:
 8:
                x_i, x_k = x_k, x_i
            end if
 9:
        end for
10:
11:
        x_i, x_{N_i-1} = x_{N_i-1}, x_i
12: end procedure
```

A runtime of O(N) can be derived, as the algorithm iterates over all particles once. Since each element in the array needs to be touched at least once to

partition the entire array there exists no better method.

Applying the algorithm to the running example looks as follows:

x	\mathbf{y}	id
0.4	0.3	0
0.2	0.6	1
0.8	0.9	2
0.6	0.5	3
0.3	0.8	4
0.7	0.1	5
0.9	0.3	6

The particles are then partitioned with a cut in the x axis set to 0.65 as seen in figure 5.

x	У	id
0.4	0.3	0
0.2	0.6	1
0.3	0.8	4
0.6	0.5	3
0.8	0.9	2
0.7	0.1	5
0.9	0.3	6

Finally $cell_2$ is cut divided into $cell_4$ and $cell_5$ with the cut position 0.55 along the y axis as seen in figure 5 ending up with:

x	\mathbf{y}	id
0.4	0.3	0
0.6	0.5	3
0.3	0.8	4
0.2	0.6	1
0.8	0.9	2
0.7	0.1	5
0.9	0.3	6

Note how all particles from $cell_4$ are contained in the range of 0-1. The particles of $cell_5$ in 2-3 and finally the particles contained in the volume of $cell_3$ can be found in 4-7.

4 Theoretical Analysis

The main goal of this thesis, is to improve the runtime of the ORB algorithm by leveraging the graphics processing unit over the central processing unit. Before implementing, it makes sense to explore if and tp which degree the performance benefits could in theory manifest themselves. For this purpose a simplified model is proposed, this model is then used to compare the runtime of the CPU version against two different GPU variants. As GPU programming is very low level, the knowledge gained while developing the model also builds a solid theoretical foundation.

4.1 General Memory Model

For a consistent terminology of a computer, a general computing model, which can be applied to most modern high performance systems is established. The model reflects a single node and its hardware components, a supercomputer may have thousands of these nodes linked together where different bandwidth are seen between individual nodes. However the internodal communication and parallelization is not important in this work, because the communication and optimization strategies are handled by parts of the PKDGrav which we adopt for this research. Both the CPU and the GPU have their own memory which are connected by a data link, where the connections are bound by the memory bandwidths. The capacity of the CPU memory bandwidth is named B_{CPU} and the GPU memory bandwidth B_{GPU} . There is a separate data link between the CPU and the GPU memory, which is commonly referred to as PCI express or NVLink (for modern NVIDIA GPU's, here the simplified term I_{GC} is used. In systems with multiple CPU's there is a link between the individual CPU's which is denoted as I_{CC} . Finally the data link between GPU's is referred to as I_{GG} .



Figure 9: Illustration of the universal computing model

4.2 Supercomputers

For Piz Daint, Summit and Eiger all relevant hardware metrics were collected and compiled to table 1. Piz Daint and Eiger were chosen, because the systems can be used to test the code. Furthermore Summit is included as it is, at the time of writing this thesis, one of the most capable supercomputers in the world.

Constant	Piz Daint [CSCS, 2022]	Summit[ORNL, 2022]	Alps (Eiger)
# Nodes	5704	4608	1024
# CPUs	1	2	2
CPU Model	Intel E5-2690 v3 [Intel, 2022]	IBM POWER9	AMD EPYC 7742[AMD, 2022]
CPU Mem.	64 GB	256 GB	??
B_C	68 GB/s	$170 \; \mathrm{GB/s}$	204.8 GB/s x 2
I_{CC}	-	64 GB/s	??
Base GHZ_C	2.9 GHZ	4 GHZ	2.25 GHZ
$Max\ GHZ_C$	3.8 GHZ	4 GHZ	3.4 GHZ
# Cores	12	22	64
Architexture	Haswell	POWER9	AMD Infinity Architecture
SIMD	AVX2	(doug) VSX	AVX2
# GPUs	1	6	0
GPU Model	NVIDIA P100 [NVIDIA, 2022e]	NVIDIA V100s [?]	-
GPU Mem. Cap.	16 GB	$16 \text{ GB} \times 6$	-
B_G	732 GB/s	$900 \text{ GB/s} \times 6$	-
I_{GC}	32 GB/s	$50 \text{ GB/s} \times 6$	-
I_{GG}	-	50 GB/s	-
GPU Tflops	9,3	16.5	-
# CUDA Cores	3584	5120	-

Table 1: Datapoints of Supercomputers

4.3 Roofline Performance Model

In a first step it is determined weather the computations are bound by memory bandwidth or the actual performance of the computing unit. The most costly computation is line 6 of the cut method (figure 2) with a runtime of $O(32 \times N)$. Oftentimes when an algorithm iterates over a large dataset performing only very little calculations on its individual elements, the limiting factor is the memory. To support this claim, a roofline models for all three systems is established. A roofline model compares arithmetic intensity in flops per byte against the actual performance of the computing chip in flops and can provide insights weather an application is memory or performance bound.

4.3.1 Estimating Flops

For modern hardware, its fairly uncommon to release flops (floating point operations per second) values. Steadily evolving SIMD instruction sets result in varying performance for different implementation details, which in turn are compiled into different assembly instructions. Depending on the algorithm, implementation details and compilation flags the c++ compiler tries to compile ideal assembly instruction sets. SIMD instructions can only be used when there is a contiguous memory access, thus in some cases a poor memory layout choice may lead to a much lower flops.

For most modern CPU chip architectures AVX is the fastest SIMD instruction set available. The common AVX2 enables the processing of 8 floating point operations per instruction with a cpi (cycles per instructions) of 0.5. The *cpi* varies depending on the chip architecture, however all relevant CPU's from figure 1 support a CPI of 0.5 along with AVX2. A lower *cpi* results in a higher efficiency, as several instructions can be completed in a single cycle.

Equation 8 defines a function to estimate the number of gigaflops for a given hardware. hz is the number of cycles which can be completed within a single second by a processing unit, meanwhile nf is the number of floats which can be processed

simultaneously in a single instruction using SIMD. Finally np represents the number of processors, where a perfect parallelization is assumed, meaning 100% of the code can be parallelized.

Variable	Meaning	Unit
S	second	
cl	processor cycle	
f	floating point operation	
i	instruction	
(g)hz	(10^{12}) processor cycles per second	$(10^{12}) \text{ cl } / \text{ s}$
nf	floating point operations per instruction	f / i
cpi	cycles per instruction	cl / i
(g)flops	(10^{12}) floating point operations per second	$(10^{12}) \text{ f / s}$
np	number of processors	

Table 2: Variables to estimate gflops

$$gflops = ghz * nf * cpi^{-1} * np$$
(8)

Since peak flops benchmarks are available for NVIDIA P100 and V100s no estimations need to be made on the GPU side.

4.3.2 Estimating Arithmetic Intensity

Arithmetic intensity is measured in FLOPS per byte or the number of floating point operations which are computed per byte loaded from memory. The count left part from the cut algorithm (line 6) in figure 2 can be translated to the following isolated c++ code:

Listing 1: Counting the particles left of a cut plane

```
for (auto p= startPtr; p<endPtr; ++p) nLeft += *p < cut;
```

Where p is a C-style array which stores the particles position and nLeft stores the number of particles which are smaller than cut. The operations per loaded float can be listed as follows:

- 1. Compare particle to cut
- 2. Add result to nLeft
- 3. Increment pointer p
- 4. Compare pointer p with endPtr

Which results in a total of 4 operations. Since a single float is stored using 4 bytes, this computes to 1.0 operations per byte or an arithmetic intensity of four.

Estimating the arithmetic intensity for the GPU is a lot more complicated as it can vary a lot depending on the specific implementation details. But for now the same arithmetic intensity is assumed for both CPU and GPU.

Note that SIMD instructions do not influence the arithmetic intensity, as the number of floating point operations per byte remains the same. It just means that a set of operations is executed concurrently. What is however influenced by AVX, is the maximum number of GFLOPS which can be processed by the hardware. If AVX was ignored, the algorithm would clearly not be bound by memory, but by the much lower performance of the processing unit. For this reason it is important to consider SIMD instructions.

4.3.3 The Plots

The maximally achievable gflops for Piz Daint are computed as follows: The datapoints from figure 1 are plugged into the formula 8 for np = 2, 4 and 8.

$$3.8 \times 10^{12} cl/s \times 8f/i \times \frac{1}{0.5} i/cl \times 2 = 121.6gflops$$
 (9)

$$3.8 \times 10^{12} cl/s \times 8f/i \times \frac{1}{0.5} i/cl \times 4 = 243.2gflops$$
 (10)

$$3.8 \times 10^{12} cl/s \times 8f/i \times \frac{1}{0.5} i/cl \times 8 = 486.4g flops$$
 (11)

The results are depicted as horizontal lines in figure 10. The memory bandwidth is plotted as a line with an equivalent slope. Finally a dotted line represents the arithmetic intensity of the Count Left procedure. To interpret the roofline mode, one has to follow the dotted line starting from the bottom. Weather it intersect with line representing the maximal performance or the memory bandwidth first, gives an indication weather the program is performance or memory bound.

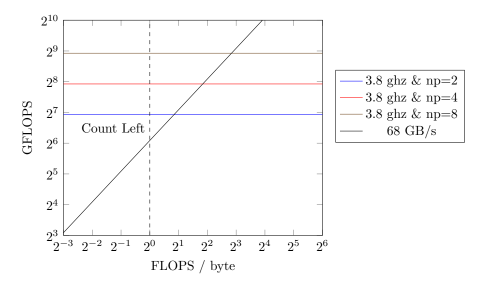


Figure 10: Roofline Model for Piz Daint CPU

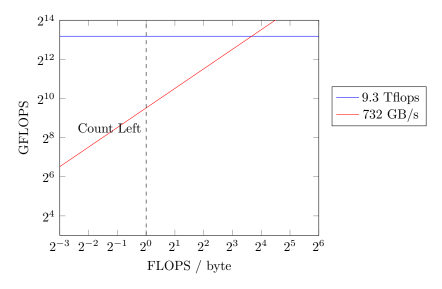


Figure 11: Roofline Model for Piz Daint GPU

As it can be seen in figure 10 and 11 both the GPU and CPU version running on Piz Daint are memory bound, but due to its much higher memory bandwidth limit, a GPU version should be favored.

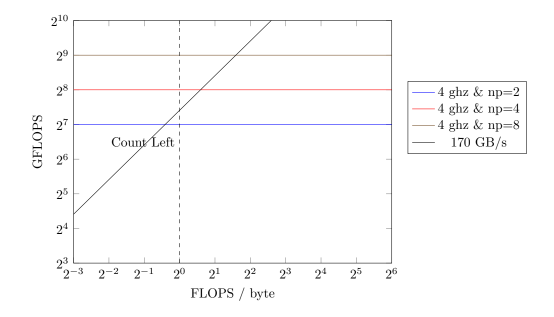


Figure 12: Roofline Model for Summit

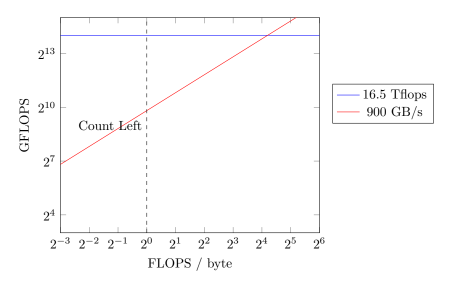


Figure 13: Roofline Model for Summit GPU

As visible in figure 12 and 13 the same applies to Summit. Its notable how the memory bandwidth only becomes the limiting factor when assuming a parallelization with 4 processors.

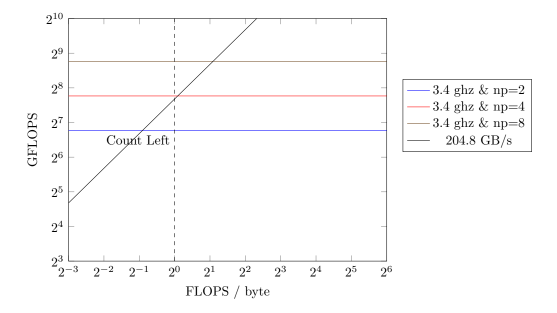


Figure 14: Roofline Model for Alps

Finally alps, which has the highest memory bandwidth compared to its flops, also becomes limited by the memory bandwidth after using more than four processors.

4.3.4 Empirical Verification

A minimal C++ code is considered to verify the theoretical model. The -S flag along with the g++ compiler is used to generate assembly code from the c++ source code. For testing purposes all entries in the array are set to random values between 0 and 1, and cut is set to 0.5.

In a first test, only O3 is enabled. The generated assembly code is depicted in listing 2.

Listing 2: Reduction Assembler Code without AVX

```
1 .L18:
2 movups (%rax), %xmm0
3 addq $16, %rax
4 cmpltps %xmm2, %xmm0
5 psubd %xmm0, %xmm1
6 cmpq %rdx, %rax
7 jne .L18
```

psubd is a packaged instruction, meaning it already uses some form of SIMD instructions. The command is used in the MME and later the SSE2 instruction set. Since the xmm registers are used, it follows that it is an AVX instruction.

If the compile flag -march = native is added, the instructions as depicted in listing 3 are the result of the compilation.

Listing 3: Reduction Assembler Code with AVX2

```
1 .L19:
2 vmovups (%rax), %ymm3
3 addq $32, %rax
4 vcmpltps %ymm2, %ymm3, %ymm0
5 vpsubd %ymm0, %ymm1, %ymm1
6 cmpq %rdx, %rax
7 jne .L19
```

 $vmovups,\ vcomplpts$ and vpsubd are AVX2 commands since they are using the ymm registers.

The following results were obtained on 2^{27} particles with a thread count of 16 on Piz Daint. The reduction takes 2.3 milliseconds, resulting in a throughput of $2^{27}/10^9 * 10^3/2.3 = 58gflops$. This is close to the memory bandwidth (68 GB/s) and far below the performance maximum of the CPU, which even for (np = 1) is 3.8*8*2*1*16 = 972.8gflops. Thus the algorithm is assumed to be memory bound for the CPU and most probably memory bound for the GPU.

4.4 Runtime Estimates

A runtime estimation for the CPU version and two different GPU implementations is constructed. In the first GPU implementation GPU Counting, an array of particles is sent to the GPU, where counting is then performed. In a more advanced variant GPU Counting and Partitioning the particles are sent from the CPU to GPU only once and the partitioning is performed on the GPU as well.

A precision p of 32 is assumed, which is general standard for both integers and floats (doug) I think things can be more complicated with float and its a sensible assumption for astrophysical simulations. The total storage required for all particle positions is $32 \times 3 \times Nbits = 4 \times 3 \times NBytes = 12 \times NBytes$. Furthermore d = 1024 and $N = 10^9$ are assumed.

4.4.1 CPU Version

Each time the leaf cells of the SPTDS are split into two child cells and appended to the tree, the tree ends up with twice the number of cells. As the goal is to construct a tree with d leaf cells, the cut algorithm needs to be performed $\lceil log_2(d) \rceil$ times over all leaf cells.

Consequently the runtime estimate sums over all $\lceil log_2(d) \rceil$ iterations, where in each iteration a cut has to be found for i cells. The cut for a single cell is found in p iterations, where the number of particles to be iterated over for a single cell is divided by two in each iteration.

$$\sum_{i=1}^{\lceil \log_2 d \rceil} i \times p \times \frac{\frac{s}{i}}{B_C} \tag{12}$$

i cancels out from the equation and the equation becomes:

$$\sum_{i=1}^{\lceil \log_2 d \rceil} p \times \frac{s}{B_C} \tag{13}$$

Which can be simplified to:

$$\lceil \log_2(d) \rceil \times \left(p \times \frac{s}{B_C} \right)$$
 (14)

4.4.2 GPU Counting

The Equation 15 for the GPU is similar, the only difference being that GPU memory bandwidth B_{GPU} is used instead of the CPU bandwidth. Furthermore the additional overhead of sending the data from the CPU to the GPU has to be reflected. This adds the terms size divided by CPU to GPU memory bandwidth denoted as I_{GC} .

$$\lceil log(d) \rceil \times \left(p \times \frac{s}{B_G} + \frac{s}{I_{GC}} \right)$$
 (15)

4.4.3 GPU Counting and Partitioning

Another variant is to performing the partitioning on the GPU, meaning there is no need to send data from the CPU to GPU each time a cut needs to be found. This allows for a reduction of the costly overheads introduced by the rather slow bandwidth I_{GC} . The overhead can be moved out of the bracket as the transfer needs to happen only once before.

$$\lceil log(d) \rceil \times \left(p \times \frac{s}{B_G} \right) + \frac{s}{I_{GC}} = t$$
 (16)

4.4.4 Plugin Values

Lets plugin the values from figure 1 into the corresponding formulas 14, 15 and 16 for Piz Daint. The naive implementation yields the following speeds for the naive CPU implementation:

$$\lceil log(1024) \rceil \times \left(32 \times \frac{12GB}{68GB/s} \right) = 56.47841s$$
 (17)

And the corresponding GPU implementation:

$$\lceil log(1024) \rceil \times \left(32 \times \frac{12GB}{732GB/s} + \frac{12GB}{32GB/s} \right) = 8.99707s$$
 (18)

This yields in a speed-up of:

$$\frac{56.47841}{8.99707} = 6.27742 \times \tag{19}$$

When considering the GPU Counting and Partitioning the results are: And the corresponding GPU implementation:

$$\lceil log(1024) \rceil \times \left(32 \times \frac{12GB}{732GB/s} \right) + \times \frac{12GB}{32GB/s} = 5.62155s$$
 (20)

This yields in a speed-up of:

$$\frac{56.47841}{5.62155} = 10.04675 \times \tag{21}$$

4.5 Conclusion

The computations for Eiger and Summit can analogously to the ones from Piz Daint. Eiger is non hybrid, thus only estimates for a CPU version are computed. All the results are plotted in 15. As expected the GPU Counting outperforms the CPU version on both hybrid (CPU and GPU available)super computers. Furthermore GPU Counting and Partitioning yields more performance improvements, but the speed up is less drastic. As the model may differ from reality depending in implementation and compilation details, these results cannot be taken for granted but offer a good baseline of what is possible.

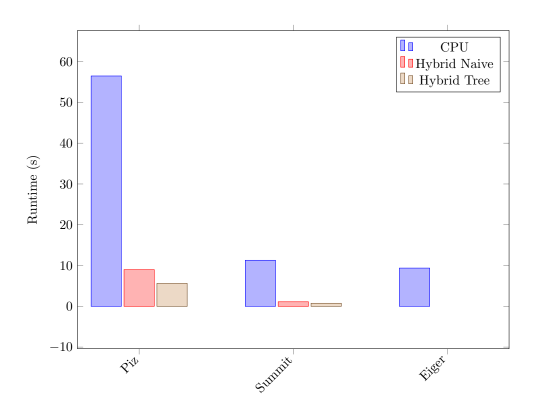


Figure 15: Execution times of different strategies

5 CPU Implementation

In this section the stand-alone implementation of the ORB algorithm using CPU acceleration only is described. Building on the CPU version, GPU accelerated alternatives to selected parts of the code are proposed.

The code is developed somewhat separately from the PKDGrav codebase in order to keep the project smaller and more easy to understand. However the machine dependent layer (MDL) from PKDGrav is used, which provides a system to distribute the workload among processors. This reduces the complexity of the implementation and communication methods and CPU parallelism are abstracted away from the code. Furthermore it makes integration into PKDGrav simple. Note that processors might be located in different nodes, but MDL is general enough to abstract node and processor level communication and parallelization away.

Not every part of the code is described in detail, however any details crucial to performance are outlined. Furthermore, special attention is paid to the CUDA kernels and memory management strategies for two reasons: minor implementation details can have a big impact upon the performance and increased replicability.

For this project C++ 17 along with CUDA 11.3, OpenMPI 4.1.0 and Blitz++ 1.0.2 [Veldhuizen, 2022] is used.

5.1 MDL and PKDGrav

MDL, the machine dependent layer provides an abstraction layer around the parallel primitives introduced by PKDGrav. The Master layer is responsible to coordinate the flow of the program. It does so by calling the PST (Processor Set Tree), which distributes the tasks and gets processors to work on them. As the name suggests, the PST is organized as a binary tree where intermediate nodes contains a pointer to a consecutive processor as well as a next lower node. Each leaf cell of the graph then correspond to an individual processor. (doug) to be precise each leaf is a "core"

Parallel processes are dispatched by descending the PST until they reach all processors. There computations are performed and the results are combined by passing them back up the PST until they reach the master. A parallel process can be programmed by implementing the service interface as explained in section 5.5.

Each processor has its own local data, which can be accessed by calling pst->lcl.

5.2 Particles

The particles array has ownership over all particle objects. It has a (N,3) shape where each row represents a single particle. To enable coalesced memory access when iterating over a single axis i.e. $x_{:,axis}$, a column major storage order is chosen. Even though row accesses patterns i.e. $x_{i,:}$ are used as well, they happen less frequently. Most importantly there are cases where entire section of a row have to copied, which is by magnitude faster with row major storage.

C-style arrays are used interchangeably with Blitz++ arrays to store all particles. Blitz++ is an open source wrapper class around C-style arrays, which helps with pointer management and can speed up the debugging process by keeping track of the array boundaries. Furthermore it features array slicing and does its own memory management. In certain cases the actual C-style array needs to be accessed, which can be accessed with array.data(). For example when iterating over the Blitz++ array,

non optimal assembly code was observed, not reflecting optimal SIMD instructions available to the hardware.

The particle data is loaded into the local data storage of each processor, meaning each processor owns a different unique set of particles.

5.3 Cell

The cell class is a structure keeping track of the fundamental cell information. In essence it is the analogue to the concept of the *cell* which has been introduced in section 3.

Quickly building and accessing elements of the SPTDS generated by ORB requires a suitable data structure. Instead of a tree with pointers, a heap can be used since the necessary conditions for a nearly complete binary tree are met. Elements within a heap can be accessed and added in constant runtime. Depicted in figure 16 is the finished SPTDS of the sample dataset stored as a heap.

The SPTDS and thus the cells are constructed on the master alone, cells are only distributed when dispatching services on the PST.

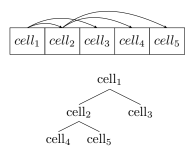


Figure 16: Tree as heap

An array can be used as an efficient storage medium for the heap, since in this case d can be set at compile time and the maximum number of cells in the SPTDS can be derived from that. Meaning the array can be allocated statically. Because MDL communicates data as arrays between the threads, another advantage of the array storage is the absence of any costly data structure conversion.

Since the STPDS is only a nearly complete binary tree and not a complete binary tree, special attention has to be paid when performing the ORB algorithm in an iterative way. But because d is known, many attributes of the SPTDS can be determined deterministically.

5.3.1 Heap Conditions

As with all heap data-structures the following conditions are given:

- The root cell has index 1
- The left child of cell at index i can be accessed at the index $i \times 2$.
- The left child of cell at index i can be accessed at the index $i \times 2 + 1$.

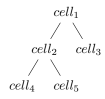


Figure 17: Tree with d=3

The following constraints can be derived given the number of leaf cells d.

- The depth of the tree can be computed with: $\lceil log_2(d) \rceil$
- $\bullet\,$ The number of leaf cells on the second last level is given by $2^{depth}-d$
- There are exactly $2 \times d 2^{depth}$ items (which must all be leaf cells) on the last level.
- The total number of cells are $2 \times d 1$.

In the example tree depicted in figure 17 the depth corresponds to: $\lceil log_2(d) \rceil = 2$, the number of leaves on the second last level is $2^2 - 3 = 1$ and the number of leaves on the last level are $2 \times 3 - 2^2 = 2$. Finally the total number of cells is $2 \times 3 - 1 = 5$.

5.3.2 Class

Among the fields in the cell are the following variables:

- *id*: Unique identification of the cell instance. Corresponds to its index in the heap array plus one. The plus one comes from the different indexing for heap constraints, starting with 1, and the classic array indexing starting with 0. It can be used to compute indices of both child cells and parent cells using the above shown formulas.
- nLeafCells: Equivalent to d_{cell} and depicts the number of leaf cells to be found in all of its successors. Its used when building the tree, to track weather a cell needs to be split or not.
- lower: 3D point coordinate representing the lower boundary corner of the 3D volume V_{cell} which is encompassed by this cells.
- upper: Represents the upper boundary corner of the volume.
- cutAxis: Axis where the cell is to be cut.

To support and simplify common operations, a further abstraction layer is given with the following functions:

- setCutAxis(): Evaluates the proper cutAxis given the upper and lower variables.
- cut(): Cuts the cell and returns two new cells.
- lower: 3D point coordinate representing the lower boundary corner of the 3D volume V_{cell} which is encompassed by this cells.
- getLeftChildId(): Returns the ID of the left child cell, meaning its volume is in the cut axis smaller than the cut plane.
- \bullet getRightChildId(): The equivalent for the right child.

5.4 Mapping Cells to Particles

As SPTDS heap is stored on the master but the particles are distributed among the local storages of the processors, its is crucial to know which particles are encompassed in the volume of which cell. Thus a data structure to keep track of the relation between cells and its particles is introduced. Thanks to the partitioning algorithm all particles belonging to a single cell can be found in a consecutive section of the particles array. A 2D array is used to keep track of the relation, which again is stored in the local data of the processor, where for each cell the corresponding range is stored as a tuple. Making the size of the array equivalent to the total number of cells $2 \times d - 1$ multiplied by two. Naturally the data structure needs to be updated across all processors whenever a partition is performed.

5.5 Services

A service implementation consists of a Service() function as well as a Combine() function. Parameters include an PST object which stores information about the process location as well as the local data. Furthermore two void pointers providing storage for the input and output data called in and out along with their respective sizes are passed. The reason void pointers are used, is the variability of the input data structure but c++ does not offer dynamic typing. Data stored inside the void pointers can be converted into their respective classes using casting.

In the Combine() function two output void pointers vout and vout2 and the sizes of the underlying arrays are given as parameters. A combination strategy of the elements can be chosen and implemented, where the results is to be stored in vout. For example in a service which returns the sum of all particles, the combine function would simply store the sum of vout and vout2 in vout.

5.5.1 Init and Finalize Service

All data stored as local data, which is not managed by Blitz++, must be allocated when initializing and after the program has finished, the memory is freed to avoid any leaks. Allocation and freeing is not free of cost, thus it makes sense to reuse memory whenever possible.

The initialize service is also responsible to load the particle data into the local memory of each PST.

5.5.2 Count Left Service

As mentioned, counting the elements smaller than the cut position is computationally intensive. T The Service() function which performs the actual computations, in points to an array of cells and as the length of the array is known, its possible to simply iterate over all of its elements and cast them to the correct cell data structure as shown in listing 4 on line 1-2. In case the foundCut flag of cell was set to true, its known that the ideal cut was already found and the cell can be skipped (line 4-6). Then the begin and end indices of the particles array corresponding to the range of objects contained within the cells volume are read and respectively a slice of the local particles array (lines 7-11) is made. Start and end pointers are computed and the cut position to be tested is fetched (lines 13 -17). Finally the number of particles smaller than the cut are counted and written back to the output array (lines 18-23). As the results for each cell needs to be stored individually, its written to the output with an offset equivalent to

the *cellPtrOffset*. This offset however does not corresponds to the *cell.id*, its rather a sequential indexing over all leaf cells of the same depth in the SPTDS.

Listing 4: Part of the Count Left Service() method

```
1 for (int cellPtrOffset=0; cellPtrOffset < nCells; ++cellPtrOffset){</pre>
      auto cell = static_cast < Cell > (*(in + cellPtrOffset));
2
      if (cell.foundCut) {
4
5
         continue;
6
      int beginInd = pst->lcl->cellToRangeMap(cell.id, 0);
      int endInd = pst->lcl->cellToRangeMap(cell.id, 1);
9
      blitz::Array<float,1> particles =
10
      pst->lcl->particlesAxis(blitz::Range(beginInd, endInd));
11
12
      float * startPtr = particles.data();
13
      float * endPtr = startPtr + (endInd - beginInd);
14
15
      int nLeft = 0;
16
      float cut = cell.getCut();
17
      for(auto p= startPtr; p<endPtr; ++p)</pre>
18
19
20
         n\, Left \; +\! = \; *p \; < \; cut \; ;
21
      out[cellPtrOffset] = nLeft;
23
24 }
```

Combining the data is straight forward, as in *Combine()* vout and vout2 contain the counts for a given number of particles. The master requires a global count across all the particles contained within all local storages, therefore each element of vout is summed together with vout2.

Listing 5: Part of the Count Left Combine() method

```
for (auto i=0; i<nCounts; ++i)
out [i] += out2[i];
```

5.5.3 Count

Counting the total number of particles contained within a cell is trivial with the cellToRangeMap. The results can be obtained by subtracting the begin index from the end index of each cell. The Combine() is equal to the one shown for the Count Left Service.

Listing 6: Part of Count Service() method

```
for (int cellPtrOffset = 0; cellPtrOffset < nCells; ++cellPtrOffset) {
    auto cell = static_cast < Cell > (*(in + cellPtrOffset));
    out[cellPtrOffset] =
    lcl -> cellToRangeMap(cell.id,1) -
    lcl -> cellToRangeMap(cell.id,0);
}
```

5.5.4 Partition

The Partitioning Service is a direct implementation of the hoare partition and can be seen in listing 7. Again the service receives a buffer with a given number of cells, which are then iterated over using the outer for loop. Each cell from the buffer is then casted to the proper struct (line 2). Using the id of the cell, the start and end indices of the corresponding particles stored in local storage can be accessed (lines 4-5). Lines 7-37 search for pairs located on the wrong side of the cut and then swap them using a simple swap() helper function. i has to be initialized using beginInd-1 as the i++ operation happens before a read on the array (line 15). For the same reason j is initialized to endInd and not endInd-1. Then the while loop terminates, i points to the first index in the array where the value is greater than the cut, therefore cellToRangeMap can be updated using i, as for the left child cell the start remains equal but the end is now i and for the right child cell its vice versa. Note that the implementation is not ideal for two reasons:

- 1. The blitz wrapper class should be avoided and instead the memory directly accessed.
- 2. In this particular case the column major storage order is not ideal, as rows are accessed over and over again.

Listing 7: Partition Service

```
for (int cellPtrOffset = 0; cellPtrOffset < nCells; ++cellPtrOffset){</pre>
      auto cell = static_cast < Cell > (*(in + cellPtrOffset));
3
      int beginInd = pst->lcl->cellToRangeMap(cell.id, 0);
4
      int endInd = pst->lcl->cellToRangeMap(cell.id, 1);
6
      int i = beginInd-1, j = endInd;
      float cut = cell.getCut();
8
      while (true)
10
11
         do
12
13
14
         } while(lcl->particles(i,cell.cutAxis)<cut and i<=endInd);</pre>
15
16
17
         do
18
19
         } while(lcl->particles(j,cell.cutAxis)>cut and j>=beginInd);
20
21
         if(i >= j) {
22
            break;
23
24
25
         swap(lcl->particles , i , j);
26
27
28
29
      swap(lcl \rightarrow particles, i, endInd -1);
30
31
      lcl->cellToRangeMap(cell.getLeftChildId(), 0) =
      lcl->cellToRangeMap(cell.id, 0);
32
      lcl->cellToRangeMap(cell.getLeftChildId(), 1) = i;
33
34
      lcl->cellToRangeMap(cell.getRightChildId(), 0) = i;
35
      lcl->cellToRangeMap(cell.getRightChildId(), 1) =
36
      lcl->cellToRangeMap(cell.id, 1);
37
38
39
```

5.5.5 Make Axis

As mentioned in section 2.1 the relevant axis to search for the cut position differs for each cell. Therefore the algorithm cannot simply iterate over a single array. In order to simplify the process the Make Axis service is introduced which iterates over all cells and copies the slices of coordinates which are encompassed the respective volumes and lie on the cells cut axis to a temporary array.

5.6 Parallel Schedule

In the context of parallelization, the number of processors is defined as np and p_0 is the processor executing the master layer. Initially it is assumed that each processor

has a random unique subset of all the particles stored in its local memory. In the running example this might look as follows:

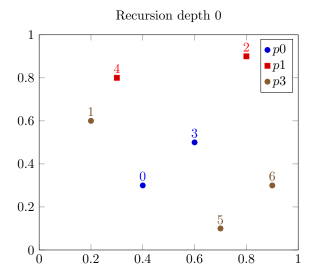


Figure 18: Example particles distributed randomly across 3 nodes

The Master schedules all other processors but also performs tasks by itself. The exact schedule is illustrated in figure 19 where a service which is dispatched from the master and executed on all processors is depicted as a horizontal rectangle. Computations which are only performed by a single processor are represented by a vertical rectangle.



Figure 19: Parallelized ORB CPU version

6 GPU Implementation

CUDA is a parallel computing platform and programming model developed by *NVIDIA*. It allows software developers and programmers to use a CUDA-enabled graphics processing unit (GPU) for general purpose processing. The CUDA platform is designed to work with programming languages such as C, C++, and Fortran and gives direct access to the GPU's virtual instruction set and parallel computational elements.

6.1 Relevant CUDA Concepts

CUDA gives us the ability to launch kernels, which are written in C with some additional CUDA specific syntax. These kernel can be run from the CPU, commonly refereed to as the host, and are then executed on the GPU, also known as the device. The device and the host have a separate memory and as seen in section 4.1 the transfer I_{CG} rates between the GPU and CPU are generally slower, under performing I_{C} and I_{G} speeds. Therefore one of key the challenges when rewriting CPU code to GPU code is to limit data transfers between the GPU and CPU as much as possible. Furthermore the problems have to be divided into smaller subproblem, such that each of them fits a single block. CUDA cannot give any guarantees considering the order of execution of these blocks greatly limiting the possibilities of communication between blocks. Algorithms have to be fundamentally redesigned when porting them from CPU to GPU code. Generally problems which are applied on large data arrays are well suited where less connection between computations is suitable the easier it is to implement an GPU version of the computation. Furthermore it can be assumed that problems with less branching work better.

The location of a thread within a block can be accessed using threadIdx.x. The block ID among all other blocks associated with the same kernel is stored in blockIdx.x. Finally the number of blocks are stored in the variable gridDim.x.

Building a tree is therefore a rather challenging problem as there are many computations which influence another and building a tree involves a lot of branching, at least when it is done in a conventional manner.

Listing 8: Calling a CUDA Kernel

```
1 add <<<
2     nBlocks ,
3     nThreads ,
4     nThreads * bytesPerThread ,
5     stream
6 >>>(
7     param1 ,
8     param2
9 );
```

A kernel is executed exactly once for every kernel, where a block consists of many kernels. The number of kernels per block and the total number of blocks can be defined by the user as seen in the kernel call syntax in figure 8 on line 2 and 3.

6.1.1 Warps

Each consecutive grouping of 32 threads form a warp. All threads within a warp are executed in parallel, given that there is no warp divergence. Because warps are executed in parallel, there is no need for synchronization between the threads of the

same warp. A warp divergence can occur, if there is a control statements, where two or more threads from the same warp execute a different code. This leads to a decrease in performance and should be avoided whenever possible.

6.1.2 Memory

Each block has access to shared memory register, where the capacity of this register can be defined at runtime by the host as seen in figure 8 on line 4. The maximum shared memory size depends depending on the architecture ranges between 48 and 96kB. [NVIDIA, 2022d] There are also upper limits for the number of threads per block which is usually between 512 and 1024. [NVIDIA, 2022d] With a shared memory size of 48kb and 1024 threads per block there is $\frac{48000B}{1024} = 46B$ of memory available per thread, which is roughly equivalent to 11 32 bit precision numbers per thread. However as the thread counts per block is usually not maxed out, there is more shared memory space available per thread.

Data in the shared memory exists only as long as the kernel exists and it cannot be accessed across different blocks. Race conditions do apply to shared memory, but threads in a block can be synchronized, therefore it is not safe to have writes from multiple threads on the same shared memory address. There exists a CUDA command __synchtreads() which allows us to synchronize all threads within a block, enabling us to control the execution order of individual statements.

Additionally to the shared memory, there exists global memory. Global memory is persistent even after a block or a kernel has finished and is the only way to send data between the device and the host. Since the data is not deleted after a kernel has finished, the data can and should be reused whenever possible. Global memory is significantly slower than shared memory and it is best practice to copy data from global memory to shared memory before performing actual computations on it, thus reducing the total number of memory accesses on global memory. After the computations have finished, the results can be written back from shared to global memory.

6.1.3 Memory Access Patterns

Global memory is fetched from memory in 32 Byte packages, which translates to $\frac{32}{4} = 8$ floats. A simple aligned sequential memory access pattern, where each tread reads a single float from global memory, results in 4 transaction per warp (32 threads). Unaligned memory access can have minor performance penalties because more memory banks have to be loaded per warp.

For shared memory each bank consists of 32 bits, which translates to a 32 bit precision number. If 32 unique threads access 32 different banks, then its considered an ideal access pattern, because the banks can be loaded in parallel. If however multiple threads access the same memory, a bank conflict is introduced and the access speed is greatly reduced.





6.1.4 Asynchronous Operations

There are two types of engines that can be used to execute kernels in CUDA streams: copy engines and kernel engines. Copy engines are used to copy data between host and device memory, and between different types of memory on the device. Kernel engines are used to execute CUDA kernels. The number of individual engines depends on the actual hardware, lower end GPUs usually have a single kernel and a copy engine, more advanced architectures can have more than one copy or kernel engine.

A CUDA stream is a sequence of commands that are executed in order on a CUDA device. Streams can be used to improve the runtime of a CUDA program by overlapping the execution of different kernels. For example, a copy kernel can be executed in one stream while a compute kernel is executed in another stream. This overlap can lead to a significant performance improvement.

6.1.5 Synchronization

Whilst threads within a block can be synchronized, this is not the case for the blocks launched from a kernel. The only real synchronization technique, is to wait for a kernel, meaning all blocks associated with this kernel, to finish and execute a consecutive task using another kernel.

Since the GPU is limited in its computing and also memory capabilities, the number of blocks which are run in parallel are limited, therefore CUDA cannot give us any guarantee, weather a set of blocks are run serially or in parallel. Some of the blocks might be run in parallel, meanwhile others are run serially. This very constraint, forces

the programmer to rethink algorithms and think of ways, how individual subproblems can be run somewhat independently of each other.

There exists however a way to communicate between blocks in a safe way. Atomic operations can be performed on global memory without any race conditions.

6.2 Streams

As described in section 6.1.4, streams can be leveraged to improve the runtime a CUDA accelerated application. In the GPU accelerated ORB streams are used in a straight forward manner, each thread performs all its computations using its own unique stream. Since leveraging streams only improves the runtime in low numbers because the actual copy and kernel engines are not very numerous, this variant is very simple and effective. It was observed that using more streams on the individual threads does not improve the runtime, quiet contrary it can have a negative impact.

6.3 Memory Management

Several strategies are used to improve the performance of memory allocation:

- Pinned Memory As the GPU cannot access the default paged memory directly, when copying memory from the host to the device, the memory must first be copied to pinned memory. This means when pinned memory is used directly with cudaMallocHost(), data transfers can be around twice as fast. Thus pinned memory is used for all data, which needs to be either sent from the host to device or vice versa.
- 2. Reuse Memory Memory allocation and freeing can be avoided altogether by reusing previously allocated memory whenever possible. Since there exists a fixed number of particles as well as an upper limit for the number of cells, memory can be allocated when calling the Initialize service and finalized with the Finalize service.

6.4 GPU Accelerated Count Left

To solve the problem on the GPU essentially a general reduction can be taken as a basis and adapted to sum elements, which fulfill a certain condition. The condition is weather an element is than a given value, which in this case is the position of the cut plane. The code is based on the reduction as its explained in a Webinar from NVIDIA. [Harris, 2022]

6.4.1 Schedule

The entire schedule of the GPU accelerated ORB is depicted in figure 20. In a first step the initialize service is called, where the necessary data is allocated on all devices and the particles are loaded. Furthermore the initial SPTDS is constructed, which essentially only consists of the root node, encompassing the entire domain and all the particles. Nextthe main loop of ORB is entered, which iterates until a SPTDS with the desired size is constructed. Within the loop the Count Service is called, computing the number of cells for each cell summed over all the threads in the system. To prepare the data for the GPU transfer a temporary array is made, using the make axis service, which is then sent to the GPU. At this stage the actual bisection methods starts on

the master, where counting the number of particles left of a cut plane is done with the GPU Count Left service. Depending on the outcome, the cut position is improved respectively and the process is repeated until a nearly perfect cut is found for all the leaf cells of the current SPTDS. For each leaf cell cell two new child cells can be generated using the computed cut positions. Finally the SPTDS can be enriched with the new cell data and the particles array is partitioned accordingly using the Partition service. The tree building loop is then repeated or if the desired size of the SPTDS is reached, the loop is exited and the Finalize service is called to free the allocated memory.



Figure 20: Parallelized ORB GPU version

6.4.2 Service

The GPU version of the Count Left Service invokes the reduction kernel exactly once for every leaf cell of the current SPTDS. Meaning on level 0 of the tree there is only one kernel per processor being executed, when proceeding to further levels, the number of kernels is equivalent to 2^{depth} per processor. The number of blocks per cell can be determined by N_{cell} divided by the number of threads per block and divided by the

number of elements per thread, which is a strategy to increase the thread occupancy.

All the relevant information can be passed along with the cell can be passed using pass by value parameters, there is no need to copy any additional data from the CPU to the GPU in advance. The only relevant data is already in global memory, when the particles where copied from the host to the device. However the results, which are also stored in global memory, need to be copied back from the GPU to the GPU after the kernel has finished.

6.4.3 Kernel Code

The actual kernel code is depicted in listing 9. The input parameters of the reduction kernel (line 13-16) are in order: the input data, which is in sequential order the particles, the output array, where the results of the kernel is stored, the position of the cut plane and finally the total number of elements contained within the cell. Because of the Make Axis service the kernel can simply iterate over a single array.

In a first step on lines 18-20 each thread ID *tid* is associated with an index in the global memory of input data. Furthermore on line 22-23 the shared data is initialized and set to zero. Its size is equivalent to the block size, the total number of thread per block. Meaning a single float can be stored per thread. On lines 25-28 the thread iterates over all elements which it is associated to and copies the results of a compare to shared memory. This allows us to work exclusively with shared memory instead of global memory, reducing the amount of costly memory accesses.

There exists an optimum in terms of number of operation performed by a single thread. In order to adapt the number of ops dynamically, the while loop ensures that each thread iterates over a certain number of elements as seen in lines 25-28. The input parameter n defines the total number of particles. Therefore if the total number of blocks is reduced by a factor of r, the while loop will iterate over r elements. Inside the while loop on line 27 the index i is incremented by gridSize in each iteration. The grid size is equivalent to the total number of threads associated with the kernel. Therefore when launching the kernel, the number of blocks can be divided by r which then results in r iterations performed in the while loop. r to 16 as more or less items per thread result in a lower performance.

The rest of the kernel code performs the actual reduction, where in each step two elements from the shared memory are summed together. Given a block size of 256, after line 37 exactly 256 sums are stored in shared memory. In a next step as seen on line 37 (lines 31-36 are not relevant as the size is smaller than 512) the left half threads in the block add their value together with the values stored in a position offset by 128. This step reduces the total number of elements to be considered to 128, and the step can be repeated as seen on lines 42 to 46.

The reduction is optimized by using a template, which indicates the total number of thread per block, which is equivalent to the blockSize. Since the template is evaluated at compile time, all the if statements taking blockSize as a parameter in figure do not cause any performance loss. In fact, because the reduction loops are unrolled (31-52) the the performance of the kernel is increased.

Each time there is a control statement involving non template arguments, a branch divergence is introduced. On the GPU these divergences become problematic bad when they are introduced inside a warp. Meaning if any of the 32 within a warp execute a different code from the other threads, the warp cannot function optimally anymore. For this reason, the *warpReduce* method as seen on line 2-9 is introduced. This method is only executed when there are 64 elements in shared memory which are

still to be summed, which requires only 32 threads as every thread in each iteration sums two elements together. Inside the warp reduce method, there are more unrolled loops continuing the same patterns as described above. However in this case there are no control statements that could introduce a warp divergence, which means there are computations performed which are not actually necessary, however as the computations inside a warp are executed in parallel anyways, this comes with no performance penalties. Inside a warp there is no need for synchronization as all threads are executed in parallel.

(doug) why don't you use the warp shuffle instrinsics, e.g., __shfl_down_sync

Listing 9: Conditional Reduction in CUDA

```
1 template <unsigned int blockSize>
  extern __device__ void warpReduce(
      volatile int *sdata, unsigned int tid) {
3
      if (blockSize >= 64) sdata[tid] += sdata[tid + 32];
     if (blockSize >= 32) sdata[tid] += sdata[tid + 16];
5
      if (blockSize >= 16) sdata[tid] += sdata[tid + 8];
6
         (blockSize >= 8) sdata[tid] += sdata[tid + 4];
     if (blockSize >= 4) sdata[tid] += sdata[tid + 2];
      if (blockSize >= 2) sdata[tid] += sdata[tid + 1];
10 }
11
  template <unsigned int blockSize>
^{12}
  extern __global__ void reduce(
13
      float *g_idata,
       unsigned int*g_odata,
15
       float cut,
16
       int n) {
17
      unsigned int tid = threadIdx.x;
18
      unsigned int i = blockIdx.x*(blockSize) + threadIdx.x;
     unsigned int gridSize = blockSize*gridDim.x;
20
21
      _shared_ int sdata[blockSize];
22
     sdata[tid] = 0;
23
24
      while (i < n) {
25
26
         sdata[tid] += (g_idata[i] <= cut);
         i += gridSize;
27
28
      _syncthreads();
29
30
      if (blockSize >= 512) {
31
         if (tid < 256) {
32
            sdata[tid] += sdata[tid + 256];
33
34
         --syncthreads();
35
36
      if (blockSize >= 256) {
37
         if (tid < 128) {
38
39
            sdata[tid] += sdata[tid + 128];
         } __syncthreads();
40
41
      if (blockSize >= 128) {
42
         if (tid < 64) {
43
            sdata[tid] += sdata[tid + 64];
44
         } __syncthreads();
```

Initializing the kernel once for each cell is a simple solution, however as the number of cells is increased, more calls to the kernel are made. Calls are not free of cost, as there is some overhead with each initialization of a kernel. Thus the performance degrades gradually as the tree is built, because on each level the number of cells for which the bisection method is executed doubles. At some point, the overhead even dominates over the actual computations by a lot.

6.5 Improved GPU Accelerated Count Left

To solve the degrading performance problem some changes are made to the kernel and the overall schedule. The main idea is to prepare the necessary data for all cells in advance, effectively reducing the overall number of necessary kernel calls to $\lceil \log_2 d \rceil$ instead of $2 \times d$. In figure 21 the resulting number of kernel calls are compared for both versions. Each block is provided with information concerning the start and end index of the cells particles, the cut position as well as an index of the cell.

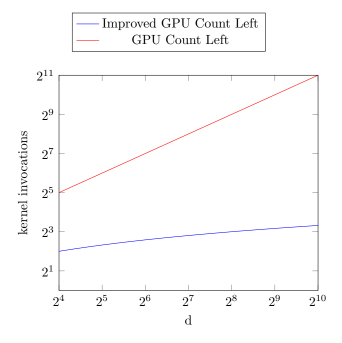


Figure 21: Kernel calls with GPU count left versions

The improved Count Left service works as follows on a simple example: Given $cell_2$

with a volume that encompasses particles in the range 0 - 10240 and $cell_3$ encompasses particles in the range 10240 - 20480. A block size of 256 threads is given and the number of elements per thread is set to 16, thus each block processes 256 * 16 = 4096elements. In this case two blocks are allocated to each cell, where the first block processes elements 0 - 4096 and the second one 4096 - 10240. By assigning more work to the second block, the threads will perform more work as others, however this is more favorable than threads which perform less work. Furthermore blocks where not even all the threads are occupied can be avoided that way. Consequently the third and fourth block respectively process particles 10240 to 14446 and 14446 to 20480. Since the number of required blocks for cell is known as well as the exact range of particles for which the reduction has to be computed, all this information can be copied to the GPU in advance and a kernel with the total block sum can be launched. Each kernel stores its information in an output array where the assignment between the output and the actual cell is known. The service therefore needs to iterate over all the output which have been computed by the blocks associated with a kernel and output them at a single cell specific location in the output ou array of the service.

6.5.1 Schedule

Only a single change is made to the schedule: A service called GPU Copy Cell Service is introduced, which does as the name says, copies the cell information from the CPU to the GPU. Since only the cut positions change while ORB is iterating to find and improve a cut and the rest of the cell information stays static, this service can be called outside of the loop. The Improved GPU Count Left Service copies the cut data each time it is invoked. Other than that, the schedule remains equal.



Figure 22: Parallelized ORB GPU version 2

6.5.2 Kernel Code

The kernel code as seen in listing 10 largely remains the same. New data pointers are passed along as a parameter where g_begin and g_end mark the begin and end of the

important particle array slice, furthermore g_cuts is an array of cell cuts for each block. The data is prepared in such a way, that each block only needs to operate over the particles contained within a single cell. blockIdx.x provides the index to read out the cell information from the input arrays (lines 20 - 22). Everything else, including the subroutine warpReduce remains equal.

Listing 10: Kernel Optimized GPU Count Left

```
1 template <unsigned int blockSize>
2 extern __global__ void reduce(
3 float * g_idata,
unsigned int * g_begins,
5 unsigned int * g_ends,
6 float * g_cuts,
7 unsigned int * g_odata) {
8    __shared__ unsigned int s_data[blockSize];
      unsigned int tid = threadIdx.x;
10
      const unsigned int begin = g_begins[blockIdx.x];
11
      const unsigned int end = g_ends[blockIdx.x];
12
      const float cut = g_cuts[blockIdx.x];
13
14
      unsigned int i = begin + tid;
15
      s_data[tid] = 0;
16
17
      // unaligned coalesced g memory access
18
      while (i < end) {
19
         s_data[tid] += (g_idata[i] <= cut);
20
         i += blockSize;
21
22
      _syncthreads();
23
24
      if (blockSize >= 512) {
25
         if (tid < 256) {
26
            s_data[tid] += s_data[tid + 256];
27
28
29
         __syncthreads();
30
31
      if (blockSize >= 256) {
         if (tid < 128) {
32
            s_data[tid] += s_data[tid + 128];
33
34
         } __syncthreads();
35
      if (blockSize >= 128) {
36
         if (tid < 64) {
37
            s_data[tid] += s_data[tid + 64];
38
         } __syncthreads();
39
40
      if (tid < 32) {
41
         warpReduce<blockSize >(s_data, tid);
42
43
      if (tid == 0) {
44
45
         g_{odata}[blockIdx.x] = s_{data}[0];
46
47 }
```

Figure 23: Reduction in CUDA

6.6 GPU Accelerated Partitioning

A GPU accelerated Count Left service is implemented successfully. Since the particles are still partitioned on the CPU, the data needs to be copied to GPU each time the SPTDS is expanded. As the Make Axis service is used to compress the axes into a single array, only N elements need to be copied, but this needs to be repeated $\lceil \log_2 d \rceil$ times. If the particles are partitioned on the GPU, it needs to be copied only once, but in this case all three coordinates have to be copied, thus $N \times 3$. Nevertheless this corresponds a fraction of the total data transfer: $\frac{3}{\lceil \log_2 d \rceil \times}$. In figure 24 its observable that a GPU partition becomes feasible, at least in terms of data transfer, after d exceeds 16.

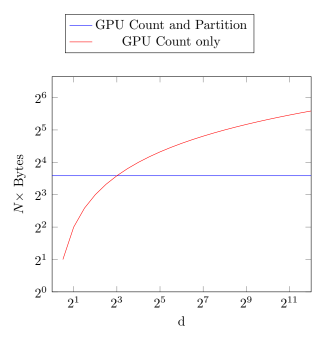


Figure 24: Data transferred with and without GPU partitioning

A GPU partitioning algorithm was first proposed by Cederman [Cederman and Tsigas, 2010] and later refined by Manca [Manca et al., 2016] where atomic operations are leveraged to improve the runtime of the algorithm. This version is built around the concepts introduced by Manca.

Early results have shown partitioning the data on the GPU is not really faster than the CPU version where the CPU version is not even implemented in a optimal manner and finally the number of particles which can be stored on the GPU memory is reduced by $\frac{5}{3}$. This is caused by required additional temporary arrays for memory swaps on the GPU. Furthermore there was not enough time to fully optimize and track down all issues with the code, therefore not full kernel code is provided, instead the underlying concepts are introduced for subsequent work.

6.6.1 Schedule

In the adapted schedule (figure 25) the Copy Particles To GPU service can be moved out of the main loop. Other than that the schedule remains the same.

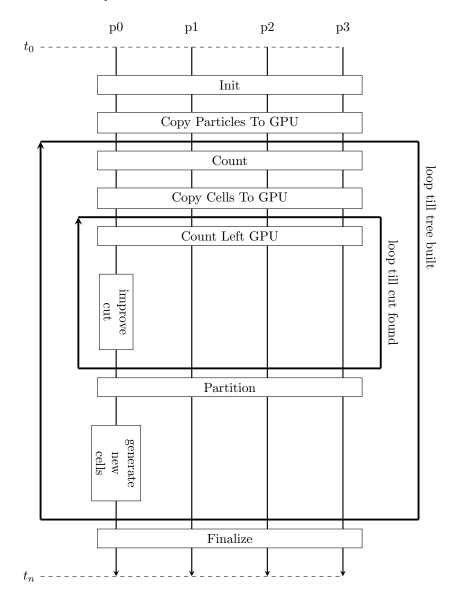


Figure 25: Parallelized ORB with GPU counting and GPU partitioning

6.6.2 Partitioning a Single Cell

In a first step a block wise partition is performed, meaning there is no synchronization between blocks at this stage. Each thread loads the particle coordinate of the matching axis into the shared memory register S. Then two copies of the data stored in the shared memory are made where the first copy A compares each value with the cut and stores 0 if the value is smaller and 1 otherwise. Consequently the second copy B stores the inverted result. Now a parallel prefix sum (also refered to as a scan) is computed on both copies, where an algorithm as described in GPU Gems [NVIDIA, 2022a] can be used. In B each element is then incremented by the an offset equal to the total number of elements smaller than the cut. In theory the block wise partition could already be applied by storing each element with index i larger than the cut at A_i and larger elements at B_i . However the goal is to partition all elements across multiple blocks. To do so a global variable called leq and g is introduced, where the total number of elements smaller equal and bigger than the cut are stored. Each time a block has computed both scans, the number of elements designated to the block smaller and larger than the cut are known. Thus a single thread of each block performs an atomic add on leq and g accordingly. AtomicAdd() returns the old value of the variable. The return value can be caught and stored two designated shared memory register s_leq and s_q . At this stage every thread in a block agrees on a value for s_leq and s_q the values can be treated as starting indices to store the block wise partitioned results in the output array. In this case however an element i, which is larger than the cut is stored location $A_i + s_l eq$ and if larger at $B_i = s_q$.

In this case the input array are coordinate values of particles and the output array is a temporary array of size N. The in an output array cannot be the same as the partition cannot be performed in place. Doing so would result in elements that are overwritten before they are read by the thread assigned to it. This cannot be avoided with the proposed technique as the relationship between the range of input data treated by a kernel and the output is not deterministic. Its dictated by the order, by which the kernel reach the atomicAdd() statement.

As of now only a single axis is partitioned, but the same needs to be done for the other two axes. One approach is to partition all the other arrays in the same kernel, as the permutations can simply be reapplied. However in this case three additional temporary arrays are needed, one for each axis. Further Reducing the number of particles which can be stored on the GPU by $\frac{1}{2}$. A more memory efficent, but slightly slower variant is to allocate another array on the GPU where the actual permutation indices are stored when the partition kernel is executed. Therefore together with the temporary array N is only reduced by $\frac{3}{5}$. After the partition kernel has finished, the coordinates can be copied back from the temporary array to the original location in a device to device copy. Then the second axis is partitioned using the precomputed permutations and the results are again copied back to the proper location. Finally after the step is repeated for the last axis, the partition is finished.

6.6.3 Partitioning Multiple Cells

The same method as introduced in section 6.5 is applied to ensure only a single kernel has to be launched for all cells. Each block is assigned a cell, where all relevant information about the cell is stored in different arrays in global memory. Each block then read the designated data by accessing the arrays at index blockIdx.x. The variables leq and g are replaced by arrays of the same length as there are cells to be partitioned

(which is not equivalent to the number of blocks). This ensures all the blocks assigned to a single cell perform the atomic operations on the same variable. For a block to know which element in the leq and g arrays it needs to access, a sequential cell index array is passed, where all blocks working on the same cell receive the same cell index.

7 Performance Analysis of ORB

All measurements where performed on Piz Daint where the performance of system can be read from table 1.

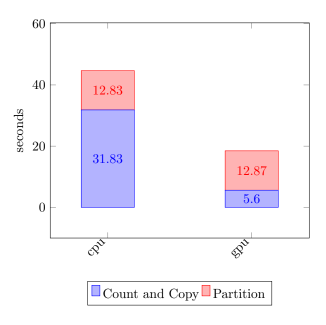


Figure 26: Execution times of different strategies

In figure 26 the final measurements are plotted. The total runtime of the ORB is around 43 seconds when performed only on the CPU, for the GPU accelerated version it comes down to 18 seconds. However a large part of the execution time can be attributed to the Partitioning, where its overhead remains equal for both versions. Even the CPU Partitioning can be improved a lot and the GPU version might increase the performance even further. If the time lost during the partitioning is subtracted, the GPU version is about 5.6 times faster, which is slightly worse than the expected speedup of 6.2. The difference can be explained by additional overheads, especially as the GPU kernel has to touch the elements more than once due to the way it was implemented.

When looking at the absolute number there is a stark difference: 56.47s predicted for the CPU version were 31.83 was the actual timing. Similarly the GPU was expected to execute ORB (without the partitioning) in 8.99s, however it only took 5.6s. The theoretical model considers an upper bound for the total number of iterations which the bisection method takes to converge. In reality this number is closer to 20 than to p which is 32 in the measured data. When considering this, the measurements are also in an absolute term relatively close to the theoretical model. $56.47s \times \frac{20}{32} = 37s$ against the actual 31.83s and $8.99s \times \frac{20}{32} = 5.99s$ compared to 5.6s.

7.1 Scaling

To make sure the GPU version scales good the total runtimes..

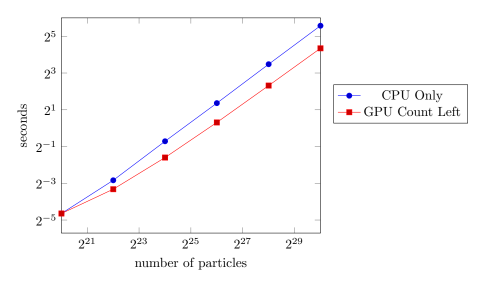


Figure 27: Execution times of different strategies

8 Conclusion

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List of Figures

Uniform random distribution of 3D coordinates in cube domain projected onto a 2d plane

10

2	Example distribution with $N = 7 \dots 1$	4
3	Example particles with ORB at recursion depth 0	4
4		4
5	Example particles with ORB at recursion depth 1	.5
6	Tree with ORB at recursion depth 1	.5
7	Example particles with ORB at recursion depth 2	6
8	Tree with ORB at recursion depth 2	6
9	Illustration of the universal computing model	22
10	Roofline Model for Piz Daint CPU	26
11	Roofline Model for Piz Daint GPU	26
12	Roofline Model for Summit	27
13		27
14	Roofline Model for Alps	28
15	Execution times of different strategies	32
16	Tree as heap	34
17	Tree with $d = 3 \dots 3$	35
18	Example particles distributed randomly across 3 nodes 4	10
19		1
20	Parallelized ORB GPU version	17
21	Kernel calls with GPU count left versions	60
22	Parallelized ORB GPU version 2	52
23	Reduction in CUDA	64
24	Data transferred with and without GPU partitioning 5	55
25		66
26	Execution times of different strategies	69
27		60
	S Commence of the commence of	
т •	. C (TD, 1, 1,	
List	of Tables	
1	Datapoints of Supercomputers	23
2		24